OM protein - nucleic search, using frame plus p2n model

Run on: December 22, 2003, 10:02:11; Search time 756.244 Seconds

(without alignments)

703.246 Million cell updates/sec

Title: US-09-446-089D-3

Perfect score: 69

Sequence: 1 KKLGYVYQDVEIP 13

Scoring table: BLOSUM62

 Xgapop
 10.0 , Xgapext
 0.5

 Ygapop
 10.0 , Ygapext
 0.5

 Fgapop
 6.0 , Fgapext
 7.0

 Delop
 6.0 , Delext
 7.0

Searched: 2888711 segs, 20454813386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seg length: 0

Maximum DB seg length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:
-MODEL=frame+ p2n.model -DEV=x1h

Q=/cgn2\_1/USPTO\_spoo1/US09446089/runat\_22122003\_100055\_23005/app\_query.fasta\_1.1 059

-DB=GenEmbl -QFMT=fastap -SUFFIX=rge -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0

-UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45

-DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000

-USER=US09446089 @CGN 1 1 5415 @runat 22122003 100055 23005 -NCPU=6 -ICPU=3

-NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG

-DEV\_TIMEOUT=120 -WARN\_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6

-FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : GenEmbl:\*

OM protein - nucleic search, using frame plus p2n model

Run on: December 22, 2003, 10:02:11; Search time 698.071 Seconds

(without alignments)

703.246 Million cell updates/sec

Title: US-09-446-089D-4

Perfect score: 56

Sequence: 1 KIVYROMVSSAK 12

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 2888711 seqs, 20454813386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+\_p2n.model -DEV=x1h

Q=/cgn2\_1/USPTO\_spoo1/US09446089/runat\_22122003\_100055\_23005/app\_query.fasta\_1.1

009

-DB=GenEmbl -QFMT=fastap -SUFFIX=rge -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0

-UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45

-DOCALIGN=200 -THR\_SCORE=pct -THR\_MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000

-USER=US09446089\_@CGN\_1\_1\_5415\_@runat\_22122003\_100055\_23005 -NCPU=6 -ICPU=3

-NO\_MMAP -LARGEQUERY -NEG\_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG

-DEV\_TIMEOUT=120 -WARN\_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6

-FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : GenEmbl:\*

OM protein - nucleic search, using frame plus p2n model

Run on: December 22, 2003, 10:02:11; Search time 1047.11 Seconds

(without alignments)

703.246 Million cell updates/sec

Title: US-09-446-089D-5

Perfect score: 101

Sequence: 1 KTPQLFFGRPYRRGDQEF 18

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 2888711 segs, 20454813386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seg length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+ p2n.model -DEV=x1h

Q=/cgn2\_1/USPTO\_spool/US09446089/runat\_22122003\_100055\_23005/app\_query.fasta\_1.1

-DB=GenEmbl -QFMT=fastap -SUFFIX=rge -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0

-UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45

-DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000

-USER=US09446089\_@CGN\_1\_1\_5415\_@runat\_22122003\_100055 23005 -NCPU=6 -ICPU=3

-NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG

-DEV TIMEOUT=120 -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6

-FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : GenEmbl:\*

OM protein - nucleic search, using frame\_plus\_p2n model

Run on: December 22, 2003, 10:02:11; Search time 1687.01 Seconds

(without alignments)

703.246 Million cell updates/sec

Title: US-09-446-089D-6

Perfect score: 140

Sequence: 1 KIDFELPXPSTTMRVRRAAHLVDDAYIXK 29

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 2888711 seqs, 20454813386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+\_p2n.model -DEV=x1h

Q=/cgn2\_1/USPTO\_spoo1/US09446089/runat\_22122003\_100055\_23005/app\_query.fasta\_1.1

- DB=GenEmbl -QFMT=fastap -SUFFIX=rge -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0

-UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR\_SCORE=pct -THR\_MAX=100 -THR\_MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -USER=US09446089 @CGN\_1 1 5415 @runat\_22122003 100055 23005 -NCPU=6 -ICPU=3

-NO MMAP -LARGEQUERY -NEG\_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG

-DEV TIMEOUT=120 -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6

-FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : GenEmbl:\*

OM protein - nucleic search, using frame\_plus\_p2n model

Run on: December 22, 2003, 10:02:11; Search time 7271.57 Seconds

(without alignments)

703.246 Million cell updates/sec

Title: US-09-446-089D-7

Perfect score: 686

Sequence: 1 RQMVSSAKTPQLFFGRPYRR.....ENAEMVRVKVRDCLDGKKLG 125

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 2888711 seqs, 20454813386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+ p2n.model -DEV=xlh

Q=/cgn2\_1/USPTO\_spool/US09446089/runat\_22122003\_100055\_23005/app\_query.fasta\_1.1

-DB=GenEmbl -QFMT=fastap -SUFFIX=rge -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0

-UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45

-DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000

-USER=US09446089 @CGN 1 1 5415 @runat 22122003 100055 23005 -NCPU=6 -ICPU=3

-NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG

-DEV TIMEOUT=120 -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6

-FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : GenEmbl:\*

FILE 'AGRICOLA, MEDLINE, BIOSIS, CAPLUS' ENTERED AT 11:04:23 ON 08 JAN 2004 15 SEA PLU=ON ((POLYPHENOL(3A) OXIDASE) OR (TYROSINASE)) AND L1 (SCROPHULARIAC? OR AGALINIS OR AGALINUS OR ALECTRA OR AMPHIANTH US OR ANGELONIA OR ANTIRRHINUM OR ASARINA OR AUREOLARIA) 10 DUP REM L1 (5 DUPLICATES REMOVED) L2 D TI 1-10 D IBIB AB 6 O SEA PLU=ON ((POLYPHENOL(3A) OXIDASE) OR (TYROSINASE)) AND L3 (BACOPA OR BARTSIA OR BELLARDIA OR BESSEYA OR BRACHYSTIGMA OR BUCHNERA OR CALCEOLARIA OR CAPRARIA OR CASTILLEJA OR CHAENORRHI NUM OR CHELONE OR CHIONOPHILA OR COLLINSIA OR CORDYLANTHUS OR CYMBALARIA) 10 SEA PLU=ON ((POLYPHENOL(3A) OXIDASE) OR (TYROSINASE)) AND T.4 (DASISTOMA OR DIASCIA OR DIGITALIS OR DIPLACUS OR DOPATRIUM OR EPIXIPHIUM OR ERINUS OR EUPHRASIA) 9 DUP REM L4 (1 DUPLICATE REMOVED) L5 D TI 1-9 D IBIB AB 9 1 SEA PLU=ON ((POLYPHENOL(3A) OXIDASE) OR (TYROSINASE)) AND L6 (FREYLINIA OR GAMBELIA OR GRATIOLA OR HEBE OR HOLMGRENANTHE OR HOWELLIELLA OR KECKIELLA OR KICKXIA OR LAGOTIS OR LATHRAEA OR LENDNERIA OR LEUCOPHYLLUM OR LEUCOSPORA OR LIMNOPHILA OR LIMOSELLA OR LINARIA OR LINDERNIA) D IBIB AB O SEA PLU=ON ((POLYPHENOL(3A) OXIDASE) OR (TYROSINASE)) AND L7 (MACRANTHERA OR MAURANDELLA OR MAURANDYA OR MAZUS OR MECARDONIA OR MELAMPYRUM OR MICRANTHEMUM OR MIMETANTHE OR MIMULUS OR MISOPATES OR MOHAVEA OR NEMESIA OR NEOGAERRHINUM OR NOTHOCHELON E OR NUTTALLANTHUS) 2 SEA PLU=ON ((POLYPHENOL(3A) OXIDASE) OR (TYROSINASE)) AND L8 (ODONTITES OR ORTHOCARPUS OR PARENTUCELLIA OR PAULOWNIA OR PEDICULARIS OR PENSTEMON OR PHYGELIUS OR PSEUDORONTIUM OR RHEMANNIA OR RHINANTHUS OR RHODOCHITON OR RUSSELIA) D TI 1-2 1 SEA PLU=ON ((POLYPHENOL(3A) OXIDASE) OR (TYROSINASE)) AND L9 (SATROCARPUS OR SCHISTOPHRAGMA OR SCHLEGELIA OR SCHWALBEA OR SCOPARIA OR SCROPHULARIA OR SELAGO OR SEYMERIA OR SIPHONOSTEGIA OR STEMODIA OR STRIGA OR SYNTHYRIS OR TONELLA OR TORENIA OR TRIPHYSARIA OR VERBASCUM OR VERONICA OR VERONICASTRUM) D IBIB AB